Molecular Phenotyping of the *pal1* and *pal2* Mutants of *Arabidopsis thaliana* Reveals Far-Reaching Consequences on Phenylpropanoid, Amino Acid, and Carbohydrate Metabolism

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The first enzyme of the phenylpropanoid pathway, Phe ammonia-lyase (PAL), is encoded by four genes in *Arabidopsis thaliana*. Whereas PAL function is well established in various plants, an insight into the functional significance of individual gene family members is lacking. We show that in the absence of clear phenotypic alterations in the Arabidopsis *pal1* and *pal2* single mutants and with limited phenotypic alterations in the *pal1 pal2* double mutant, significant modifications occur in the transcriptome and metabolome of the *pal* mutants. The disruption of PAL led to transcriptomic adaptation of components of the phenylpropanoid biosynthesis, carbohydrate metabolism, and amino acid metabolism, revealing complex interactions at the level of gene expression between these pathways. Corresponding biochemical changes included a decrease in the three major flavonol glycosides, glycosylated vanillic acid, scopolin, and two novel feruloyl malates coupled to coniferyl alcohol. Moreover, Phe overaccumulated in the double mutant, and the levels of many other amino acids were significantly imbalanced. The lignin content was significantly reduced, and the syringyl/guaiacyl ratio of lignin monomers had increased. Together, from the molecular phenotype, common and specific functions of PAL1 and PAL2 are delineated, and PAL1 is qualified as being more important for the generation of phenylpropanoids.

INTRODUCTION

The phenylpropanoid compounds derived from the shikimic acid pathway are precursors to diverse classes of phenolics, such as lignin, flavonoids, isoflavonoids, coumarins, and stilbenes. One of the intriguing characteristics of phenylpropanoids is the rapid modification in fluxes and rates of their synthesis and metabolism. This aspect is corroborated by the occurrence of specific phenolics in certain cell types, in response to specific stimuli (light, pathogens, and wounding) or during particular developmental stages.

The end products of the shikimate pathway, the aromatic amino acids Phe, Trp, and Tyr, are generated from the precursors

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phospho-enol-pyruvate and erythrose-4-phosphate. Phe ammonia-lyase (PAL; EC 4.3.1.5), the first enzyme of the phenylpropanoid pathway, removes Phe from aromatic amino acid biosynthesis to generate trans-cinnamic acid and NH₃ in a nonoxidative deamination. Therefore, PAL commits the flux of primary metabolites into the phenylpropanoid pathway and becomes rate limiting when reduced below a threshold of 20 to 25% in tobacco (Nicotiana tabacum) (Bate et al., 1994; Sewalt et al., 1997). Many studies have revealed the regulatory mechanisms for various PAL isoforms in response to different stimuli. PAL activity is mainly regulated via de novo synthesis of the enzyme, but it can also be modulated by inactivation of the protein. PAL transcript levels increase when furnished with its substrate Phe as, among others, evidenced through feeding studies in lignifying Pinus taeda cells (Anterola et al., 2002). PAL activity and PAL gene transcription are inhibited by transcinnamic acid, its product (Bolwell et al., 1986; Mavandad et al., 1990; Blount et al., 2000). Moreover, PAL is subject to posttranslational phosphorylation by a specific calcium-dependent protein kinase (Cheng et al., 2001).

Given that alterations in the expression of monolignol biosynthesis genes can lead to differential accumulation of pathway intermediates that, in turn, can affect the transcription of other

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genes (Boerjan et al., 2003), we hypothesize that it would be possible to reveal the full spectrum of the PAL function and to dissect functional redundancy of PAL isoforms by means of thorough molecular phenotyping. We have recently identified the 34 *Arabidopsis thaliana* genes that encode the 10 known enzyme functions in monolignol biosynthesis (Raes et al., 2003). The Arabidopsis genome harbors four *PAL* genes that fall into two phylogenetic groups consisting of *PAL1* and *PAL2*, and *PAL3* and *PAL4*. On the basis of their strong expression in the inflorescence stem, phylogenetic relationship, and occurrence of promoter elements, *PAL1* and *PAL2* were selected as good candidates for monolignol production during developmental lignification of vascular tissues (Raes et al., 2003).

Here, we show that in the absence of phenotypic alterations in the pal1 and pal2 single mutants and with limited phenotypic alterations in the pal1 pal2 double mutant, significant modifications occur at the transcriptome and metabolome levels in these mutants. Transcript-profiling approaches reveal far-reaching consequences of the disruption of the PAL function on the transcription of genes that encode enzymes of the phenylpropanoid, carbohydrate, and amino acid metabolisms, establishing a clear link between primary and secondary metabolisms. Furthermore, many of these alterations at the transcriptional level are translated into the corresponding biochemical effects as witnessed by targeted metabolite profiling. Our data indicate common as well as discrete functions of PAL1 and PAL2 and demonstrate that a single mutation can profoundly affect the transcription of a plethora of genes, also outside the phenylpropanoid pathway, and the biosynthesis of a variety of metabolites.

RESULTS

Morphological Characterization of the pal1, pal2, and pal1 pal2 Mutants

T-DNA insertion mutants for *PAL1* and *PAL2* were generated by exon-trap mutagenesis in the Arabidopsis ecotype C24 (Babiychuk et al., 1997; see Methods). T-DNA insertions resided in the first intron of both genes (data not shown).

Although flavonoid deficiency is known to influence several aspects of plant development, such as lateral root formation, inflorescence growth, and seed viability, the pal1 and pal2 exontrap mutants had no visible phenotypic alterations. The pal1 pal2 double mutant exhibited no gross morphological differences either, except that it had become sterile. As shown in Figure 1, the siliques remained small (3 to 4 mm versus ~12 mm in the wild type). The empty ovule sacs had initially enlarged but shriveled later, suggesting male sterility. Reduced pollen viability was also observed in transgenic tobacco downregulated for PAL (Elkind et al., 1990). Because of the failure to produce viable seeds, the double mutant generated recurrent flushes of accessory paraclades that also failed to produce seeds. These flushes continued long after the wild type and single mutants had ceased growth.

Although no growth differences could be seen with respect to time of bolting, height, or diameter of the inflorescence stem, slight differences in lignin accumulation were observed by lignin histochemistry in inflorescence stems of 2.5-month-old plants. As shown in Figure 1, a small reduction in lignin deposition was detected in the pal1 mutant and a more significant reduction in the pal1 pal2 double mutant (Figures 1A to 1L). Cell walls were further investigated at the ultrastructural level by transmission electron microscopy, as presented in Figure 2. The cell walls of xylem and interfascicular fibers were evenly thickened and uniformly stained in the wild type (Figures 2A and 2E). Double mutants were indistinguishable from the wild type in terms of cell size, cell number, or thickness of the cell wall. However, the cell wall of the interfascicular fibers and the large vessels appeared unevenly stained, and cell corners were often whiter (Figures 2B and 2F). In several cells, various signs of a reduced cell wall strength were observed: layers of the cell wall were loosened and had numerous little holes, or cellulose microfibrils became apparent (Figures 2C, 2D, 2G, and 2H). The latter phenomena were never observed in the wild type. At a low frequency, cell walls of the double mutant were aberrantly shaped and irregular (Figure 2B), similar to what has been observed in a more dramatic way in the lignin-deficient irregular xylem 4 Arabidopsis mutants (Jones et al., 2001).

In conclusion, *pal1* and *pal2* mutants have no obvious visible phenotype. Only the combination of both mutations in the double mutant leads to infertility and clear changes in lignin accumulation and secondary cell wall ultrastructure.

Expression of the Four Arabidopsis *PAL* Genes Is Altered in the Mutants

One possibility to explain the limited phenotypic differences in the pal single and double mutants is that the disruption of the PAL1 and PAL2 function is compensated for by other PAL isoforms. A comprehensive screen of the complete Arabidopsis genome revealed four PAL genes (Raes et al., 2003). The PAL genes are expressed in many tissues (including seed, seedling, leaf, root, flower, silique, and inflorescence stem), with PAL3 being generally expressed at a lower level (Wanner et al., 1995; Raes et al., 2003). Expression of PAL1 and PAL2 is detected most abundantly in roots and stems, where it increases during the later stages of inflorescence stem development. Because PAL1, PAL2, and PAL4 genes are most abundantly expressed during the later stages of inflorescence stem development (Raes et al., 2003), major effects of knock downs should manifest themselves in the inflorescence stem. Semiguantitative RT-PCR with specific primers was performed to clarify to what extent the expression levels of each of the four genes were affected in the mutants. As shown in Figure 3, T-DNA insertions in the PAL1 and PAL2 genes effectively disrupted the expression of these genes in the respective mutants. On the other hand, transcript levels of the PAL1 and PAL2 genes were increased in the pal2 and pal1 mutants, respectively, indicating a compensatory response (Figure 3). Also, the PAL4 gene was significantly induced in all mutants. The impact of PAL3, although altered in the pal2 mutant, was presumably weak because of its generally low expression level (Figure 3; Wanner et al., 1995). Therefore, the PAL transcripts are mainly derived from PAL2 and PAL4 in pal1 mutants, from PAL1 and PAL4 in pal2 mutants, and from PAL4 in the double mutant.

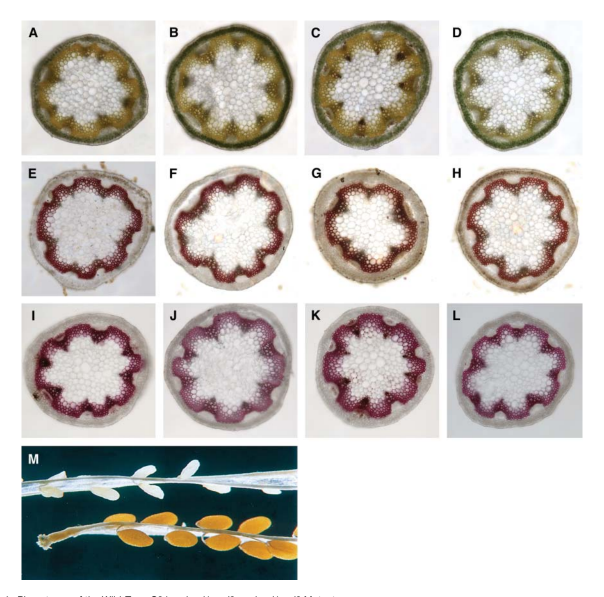


Figure 1. Phenotypes of the Wild-Type C24 and pal1, pal2, and pal1 pal2 Mutants.

Lignin staining in the basal part of 3-month-old inflorescence stems ([A] to [L]) and silique phenotype (M).

- (A) The wild type stained with aniline sulfate.
- (B) pal1 mutant stained with aniline sulfate.
- (C) pal2 mutant stained with aniline sulfate.
- (D) pal1 pal2 mutant stained with aniline sulfate.
- (E) The wild type stained with KMnO₄:HCl:NH₃.
- (F) pal1 mutant stained with KMnO₄:HCl:NH₃.
- (G) pal2 mutant stained with KMnO₄:HCl:NH₃.
- (H) pal1 pal2 mutant stained with KMnO₄:HCl:NH₃.
- (I) The wild type stained with phloroglucinol:HCl.
- (J) pal1 mutant stained with phloroglucinol:HCl.
- (K) pal2 mutant stained with phloroglucinol:HCl.
- (L) pal1 pal2 mutant stained with phloroglucinol:HCl.
- (M) Siliques of the wild type (bottom) and pal1 pal2 mutant (top) at seed maturity of the wild type.

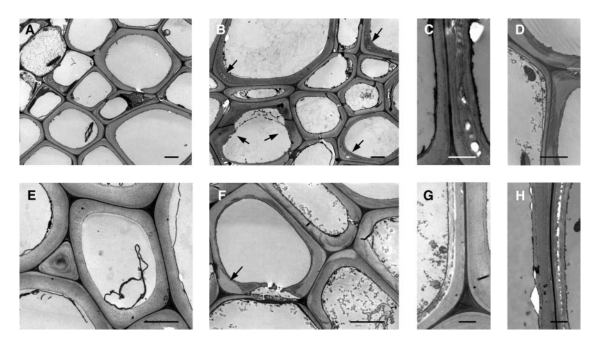


Figure 2. Cell Wall Ultrastructure in the Basal Part of 3-Month-Old Inflorescence Stems of the Wild Type and pal1 pal2 Mutants.

- (A) Xylem region of the wild type.
- (B) Xylem region of pal1 pal2 mutants. Arrows point to aberrantly shaped walls and to regions punctured with little holes.
- (C) and (D) Cell walls in the xylem region of pal1 pal2 mutants. In (D), loosened cellulose fibrils become apparent.
- (E) Interfascicular fiber region of the wild type.
- (F) Interfascicular fiber region of pal1 pal2 mutants. Arrow points to whiter cell corners.
- (G) and (H) Cell walls in the interfascicular fiber region of pal1 pal2 mutants.
- Bar in (A), (B), (E), and (F) = 2.5 μ m; bar in (C), (D), (G), and (H) = 1 μ m.

To estimate the consequences of upregulation of other isoforms at the enzyme level, PAL activity was determined in inflorescence stems of greenhouse-grown plants. As shown in Figure 4, PAL activity in the inflorescence stem was clearly reduced in the *pal1* but not in the *pal2* mutant. In the double mutant, the 25% residual PAL activity must be derived mainly from *PAL4* transcripts (Figures 3 and 4). Additional activity measurements in siliques and leaves of these plants revealed none and a 50% reduced activity in all genotypes, respectively (data not shown). Thus, most clear-cut differences in activity

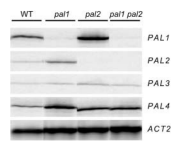


Figure 3. Expression of *PAL* Genes in 10-Week-Old Inflorescence Stems of the Wild Type and *pal1*, *pal2*, and *pal1 pal2* Mutants.

Semiquantitative RT-PCRs were performed on a pool of 10 individuals of each respective genotype (see Methods). The experiment was repeated with similar results.

were observed in the inflorescence stem, the tissue where *PAL1* and *PAL2* are most highly expressed. However, the lesion in *PAL2* was almost completely compensated for at the enzyme level. Following the PAL activity during inflorescence growth, PAL activity was highest in 10-week-old, seed-setting but still green plants (Figure 4; data not shown). Assuming that at this age the differences in PAL activity between the wild type and mutants would also be highest, inflorescence stem material of 10-week-old plants was used to conduct all further experiments.

Expression of 10 Genes Encoding Enzymes of Monolignol Biosynthesis Is Strongly Altered in the *pal* Mutants

Our search was extended for consequences of PAL disruption on the transcription of the remaining 30 genes encoding the nine other enzymes of monolignol biosynthesis (Raes et al., 2003). Their expression was assayed by RT-PCR with gene-specific primers for each of the genes. Given the rapid alterations in expression levels known to occur for many phenylpropanoid genes and the semiquantitative nature of RT-PCR, a conservative approach of fourfold change was taken to delineate those genes with clear changes in expression. As shown in Figure 5, 23 of the 30 genes were expressed in 10-week-old inflorescence stems. Among the genes that were judged important for the developmental lignification of vascular tissues based on expression, phylogenetic relationship, and occurrence of promoter elements (Raes et al., 2003), pronounced changes in expression were

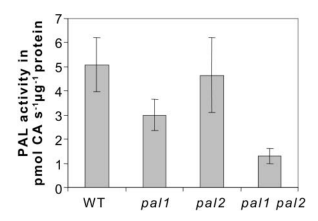


Figure 4. PAL Activity in 10-Week-Old Inflorescence Stems of the Wild Type and *pal1*, *pal2*, and *pal1* pal2 Mutants.

PAL activity of crude extracts was determined in inflorescence stems (n = 10) and is expressed as average PAL activity in pmol cinnamic acid (CA) s⁻¹ μ g⁻¹ protein with standard deviation.

found for 4-coumarate:CoA ligase (4CL1), para-coumarate 3-hydroxylase (C3H1), hydroxycinnamoyl-CoA:shikimate/quinate hydroxycinnamoyltransferase (HCT), caffeic acid O-methyltransferase (COMT), and cinnamyl alcohol dehydrogenase (CAD6). Interestingly, similar trends in alteration of gene expression were observed for most of them in both single pal mutants (Figure 5). Only the expression of C3H1 was higher in the mutants, whereas that of HCT, COMT, and CAD6 was lower (Figure 5). C3H1, hydroxylating the C3 position of p-coumaroyl shikimic/quinic acid, might be very prone to transcriptional regulation by the concentration of certain pathway intermediates. On the other hand, HCT activity is positively regulated by cinnamic acid (Lamb, 1977); hence, the decreased transcription probably correlated with the reduced synthesis of cinnamic acid in the pal mutants. Moreover, the HCT promoter contains both an H and a G box, known to be necessary for feed-forward induction by p-coumaric acid (Loake et al., 1992; Lindsay et al., 2002; Raes et al., 2003). Thus, our data underscore the responsiveness of the HCT gene to regulation by early pathway intermediates.

Caffeoyl-CoA 3-O-methyltransferases CCoAOMT5, CCoAOMT6, and CCoAOMT7, CAD2, and cinnamoyl-CoA reductase (CCR2) with an altered expression in pal mutants had not been selected as candidates for a role in developmental lignification (Figure 5; Raes et al., 2003). CCR2, assigned a primary function in pathogen response (Lauvergeat et al., 2001), was also induced in the cellulose-deficient mutants detiolated 3, radially swollen 1, and ectopic lignification 1-1 (Caño-Delgado et al., 2003). Thus, CCR2 as well as the others might become subject to transcriptional regulation upon alterations in the concentration of pathway intermediates.

pal Mutants Are Affected in Transcription of Genes Encoding Enzymes of Amino Acid, Phenylpropanoid, and Carbohydrate Metabolisms

In view of the dramatic alterations in transcription of the most closely related genes, we investigated the consequence of the pal mutations on the transcriptome by a near genome-wide cDNA-amplified fragment length polymorphism (AFLP) transcript profiling of 10-week-old inflorescence stems. Transcript profiling was performed on two biologically independent sets of the wild type and the three mutants. Each single sample was a pool of 10 individual plants to normalize for the expected variability in gene expression of individual plants. In the expression profiles, differential transcript-derived fragments were identified by their increase/decrease in one or all mutants as compared with the wild type. Fragments were withheld only when a similar expression pattern was observed in both biological sets. For 63 differential sequence tags, selected in this manner and classified in Table 1 according to functional categories, the corresponding genes in Arabidopsis were identified. Among these 63 genes, three groups accounted for one-third of the sequences (or 43% of the genes with a known function): genes related to amino acid synthesis and metabolism, to the phenylpropanoid pathway, and to carbohydrate metabolism (Table 1).

The transcriptomes of pal1 and pal2 mutants from the same biological material were also compared with the wild type with a 6K-cDNA microarray. As shown in Table 2, expression significantly changed (twofold at P = 0.001) for 36 genes, of which only three differential genes were common in both mutants. Convincingly, half of these genes (or 61% of the genes with a known function) fell into the same three functional categories (amino acid, phenylpropanoid, and carbohydrates) as in the cDNA-AFLP experiment. Only two genes were found with both methods: a homolog of Asn synthase (ASN3, At3g15450) and a putative protein (At1q73330). The fact that 27 of the 63 tags found in the cDNA-AFLP experiment are present on the microarray suggests that many of these tags were not detected because of crosshybridization with transcripts of other gene family members or transcript abundance. Clearly, cDNA-AFLP detected more differentially expressed genes involved in signal transduction/gene regulation and unknowns than the microarray (15 versus 3 and 14 versus 5, respectively). This observation underscores the complementarity of both approaches and the higher sensitivity of the cDNA-AFLP method for low-abundant messengers and members of gene families.

Figure 6 shows an interesting trend by clustering all 97 tags according to their expression in the mutants: the largest group of genes linked specifically to the pal1 mutation identifies genes of the phenylpropanoid pathway. Remarkably, $\sim 50\%$ of the differentials found in pal1 and pal2 mutants were common to pal1 and pal2, and 27 of 32 of these common differentials were also found in the double mutant (Figure 6). Only one gene (At3g12300, unknown protein; Table 1) required both mutations for a change in expression, altogether corroborating that the differentially expressed genes were linked to the defect in PAL1 and PAL2, and not to stochastic gene expression.

As summarized in Figure 7, 43 and 61% of the differentially expressed genes with a known function identified through cDNA-AFLP and microarray, respectively, were related to amino acid, phenylpropanoid, and carbohydrate metabolisms. With respect to amino acid metabolism, three tags recognized enzymes of the aromatic amino acid biosynthesis and metabolism: chorismate mutase (CM), the first committed step in Phe and Tyr biosynthesis; Trp synthase (TSB), performing the final step in biosynthesis of Trp; and Tyr transaminase (TAT), degrading Phe

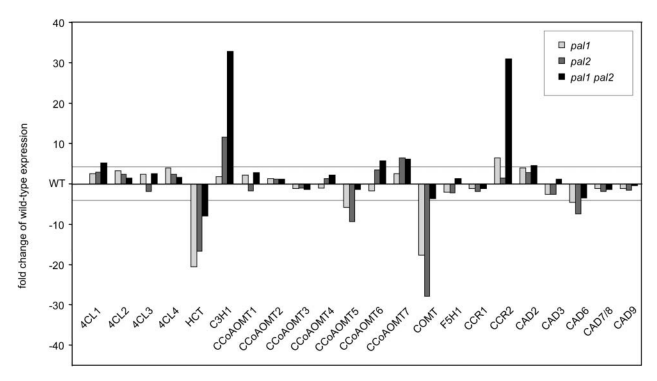


Figure 5. Expression of the Monolignol Biosynthesis Genes in pal Mutants.

Genes (34) were annotated with sensitive annotation methods for the 10 enzymes currently known to participate in the generation of monolignols (Raes et al., 2003). Semiquantitative RT-PCRs were performed on two pools of 10 inflorescence stems of each respective genotype (see Methods). Expression levels were normalized to *ACT2* gene expression before calculation of fold changes in expression levels taking the wild-type level as reference. Bars represent the average expression level of two biological repeats. Lines mark the level of fourfold upregulation or downregulation. Note that the following genes were not found to be expressed at this stage of inflorescence stem development (10-week-old plants): *C4H*, *C3H2*, *C3H3*, *F5H2*, *CAD1*, *CAD4*, and *CAD5*. For primers and gene nomenclature, see Raes et al. (2003).

and Tyr (Tables 1 and 2, Figure 7). Two genes that are homologs of ASN3 are upregulated in pal2 mutants (Tables 1 and 2). Four other tags correspond to genes encoding components of the Gly cleavage complex, primarily involved in photorespiration: two different homologs of Gly dehydrogenase and two DNA binding proteins (At1g69570 and At3g25990) that recognize motifs present in the H protein promoter with the H protein itself being part of the Gly decarboxylase enzyme complex (Tables 1 and 2). In addition, Met synthase, upregulated in the pal2 mutant, represents an interesting tag because it catalyzes the methylation of homocysteine with 5-methyl-tetrahydrofolate as methyl donor. This reaction does not only occur during de novo synthesis but also during recycling of S-adenosyl-L-homocysteine after S-adenosyl-Met (SAM) has donated its methyl group. Besides the incorporation of Met into proteins, 80% of Met is converted into SAM for various transmethylations that are also very frequent in the phenylpropanoid pathway (Figure 7).

Among the phenylpropanoid pathway-related transcripts, PAL1, PAL2, and 4CL1 were expressed in a manner consistent with the results from the RT-PCR experiment (Tables 1 and 2, Figures 3 and 5). The NADPH ferrihemoprotein reductase ATR3, found particularly induced in the pal1 mutant, is probably part of a microsomal P450 system that associates multiple P450 enzymes and NADPH:cytochrome P450 reductases. ATR homologs from poplar ($Populus\ trichocarpa\ \times\ P.\ deltoides$) and

Arabidopsis were shown to efficiently support the activity of the P450 enzyme cinnamate 4-hydroxylase (C4H), when coexpressed in yeast (Urban et al., 1997; Mizutani and Ohta, 1998; Ro et al., 2002). Differential tags from a carboxymethylenebutenolidase, dihydroflavonol 4-reductase (DFR), sinapoylglucose: malate sinapoyltransferase (SNG1), a UDP-glucosyl transferase, a glutathione S-transferase (GST), and a glutathione S-conjugate pump (GS-X pump) correspond to enzymes active toward the endpoints of biosynthetic pathways originating from the phenylpropanoid pathway (Figure 7). The UDP-glucosyl transferase belongs to the glycosyltransferase family 1; characterized members use phenylpropanoids, such as flavonol and sinapate, as substrates (http://afmb.cnrs-mrs.fr/CAZY/). The GSTs of the ϕ and τ subgroups, to which the downregulated GST also belongs (GSTF12; Dixon et al., 2002), have been implicated in cytoplasmic stabilization of flavonoids and flavonoid conjugation before uptake into the vacuole (Mueller et al., 2000; Dixon et al., 2002). The ABC transporter (AtMRP2) identified here encodes a functional GS-X pump of the plant vacuolar membrane with a high capacity for glutathionated anthocyanins (Lu et al., 1998; Liu et al., 2001; Sánchez-Fernández et al., 2001).

Perturbations in the PAL function seem to lead to adjustments in carbon fixation and carbohydrate metabolism (Tables 1 and 2, Figure 7). Ribulose-1,5-bisphosphate carboxylase/oxygenase (Rubisco) is upregulated in *pal1* and *pal2* mutants, and Rubisco

Table 1. Identities and Expression Patterns of the Sequence Tags Isolated for Their Differential Expression in the Wild Type and *pal* Mutants by cDNA-AFLP Transcript Profiling

	Expr	ession ir	1			Nearest Homo		
AGI	WT	pal1	pal2	DM	Description (MIPS)	AGI/GI	Description	E Value
mino Acid-	Related	d						
t3g04940	0	0	1	0	Cys synthase (AtcysD1)			
t5g17920	0	0	1	0	Met synthase			
xt1g69570	0	0	1	1	H-protein promoter binding factor-2b, putative	At5g39660	Promoter-binding protein like	1×10^{-118}
xt3g15450	0	0	1	1	Unknown protein	At5g10240	Asn synthetase ASN3	$3 imes 10^{-71}$
t4g33010	0	1	0	0	P-Protein-like protein	GI:20471026	Gly dehydrogenase, Homo sapiens	0
xt2g26080	0	1	1	1	Putative Gly dehydrogenase	GI:20471026	Gly dehydrogenase, <i>H. sapiens</i>	0
At2g27820	0	1	1	1	Putative chorismate mutase/ prephenate dehydratase		·	
\t5g54810	0	1	1	1	Trp synthase β chain 1 precursor			
Carbohydrat	e-Relat	ed						
At2g21170	0	0	1	0	Putative triosephosphate isomerase			
At1g32900	0	1	0	0	Granule-bound starch synthase-like protein			
At5g17420	0	1	0	1	Cellulose synthase catalytic subunit (IRX3)			
At1g35580	0	1	1	1	Invertase, putative			
xt3g48530	0	1	1	1	Unknown protein	GI:7300717	SNF4Aγ gene product, <i>Drosophila melanogaster</i>	5×10^{-64}
t5g58090	0	1	1	1	β-1,3-glucanase 6		,	
t3g46970	1	0	1	1	Starch phosphorylase H (cytosolic form)-like protein			
henylpropa	noid-R	elated						
\t3g53260	0	1	0	0	Phe ammonia-lyase (AtPAL2)			
xt3g02280	0	1	0	0	Putative NADPH-ferrihemoprotein reductase ATR3	At4g24520	NADPH-ferrihemoprotein reductase ATR1	0
At1g51680	0	1	0	1	4-Coumarate:CoA ligase (At4CL1)	At4g05160	4-Coumarate:CoA ligase-like protein	1×10^{-147}
At1g43620	0	1	1	1	Hypothetical protein	GI:6562348	UDP-glucoronosyl/glucosyl transferase, Caenorhabditis elegans	1 × 10 ⁻¹²⁴
At2g32520	0	1	1	1	Putative carboxymethylene- butenolidase		5.6gae	
At1g49140	1	0	0	0	Similar to NADH-ubiquinone oxidoreductase precursor			
_ight-Related	d							
At2g43280	0	0	1	0	Unknown protein	At5g28530	Far-red-impaired response protein (FAR1)-like	3×10^{-52}
\t1g65070	0	1	1	1	(No name given)	GI:33876770	mutS protein homolog 5, H. sapiens	1×10^{-150}
xt1g53090	1	1	0	1	Phytochrome A suppressor SPA1, putative	At4g11110	COP1-like protein	2×10^{-71}
xt3g22170	1	1	0	1	Far-red-impaired response protein, putative	At5g28530	Far-red-impaired response protein (FAR1)-like	1×10^{-141}
Stroop Doors	2002				•		•	
Stress Respo Nt3g11170	onse 0	1	1	1	ω-3 Fatty acid desaturase,			
\t4g16070	0	1	1	1	chloroplast precursor Unknown protein	At3g49050	Calmodulin binding heat shock-like protein	9×10^{-61}

(Continued)

Table 1. (co	ontinue	ed)						
	Expr	ession	in			Nearest Homo		
AGI	WT	pal1	pal2	DM	Description (MIPS)	AGI/GI	Description	E Value
At4g16860	0	1	1	1	Disease resistance RPP5- like protein			
At5g42650	0	1	1	1	Allene oxide synthase			
At5g62130	0	1	1	1	Unknown protein	GI:83226	PER1, Saccharomyces cerevisiae	1×10^{-99}
Signal Trans	ductio	n/Trans	cription	al Reg	ulation			
At3g11500	0	0	1	0	Putative small nuclear ribonucleoprotein polypeptide G			
At1g07410	0	0	1	1	Small G protein, putative			
At5g57210	0	0	1	1	Microtubule-associated protein-like	GI:12188746	Rab6 GTPase-activating protein, <i>H. sapiens</i>	4×10^{-50}
At1g04390	0	1	0	0	Hypothetical protein	GI:6705975	KIAA0740 protein, H. sapiens	2×10^{-54}
At1g21920	0	1	0	1	Phosphatidylinositol-4-phosphate	GI.0703973	Nimaurau protein, rr. sapiens	2 ^ 10
At1g23230	0	1	1	0	5-kinase-like protein Conserved hypothetical	GI:7291500	CG3695 gene product,	0
At1g23380	0	1	1	1	protein Knotted-like homeobox	At5g25220	D. melanogaster KNAT3 homeodomain protein	1×10^{-44}
At3g28450	0	1	1	1	protein, putative Receptor kinase, putative	At5g16000	Receptor protein kinase-like	1×10^{-117}
At3g61570	0	1	1	1	Putative protein	GI:29387238	protein CDC42 binding protein kinase	2 × 10 ⁻²⁹
At4g18010	0	1	1	1	Putative inositol polyphosphate	GI:21396493	β, <i>H. sapiens</i> Inositol polyphosphate	1×10^{-101}
					5-phosphatase At5P2		5-phosphatase, H. sapiens	
At5g14670	0	1	1	1	ADP-ribosylation factor-like protein			64
At2g41090	1	0	0	0	Calcium binding protein (CaBP-22)	At4g04710	Putative calcium-dependent protein kinase	5×10^{-34}
At1g73080	1	0	1	1	Unknown protein	At5g07280	Receptor-like protein kinase-like protein	0
At2g21660	1	0	1	1	Gly-rich RNA binding protein			
At3g12300	1	1	1	0	Unknown protein	GI:7297748	CG17118 gene product, D. melanogaster	5×10^{-48}
Transport								
At5g03555	0	1	0	0	Uracil transporter-like protein			
At5g02270	0	1	0	1	ABC transporter-like protein			
At4g34950	0	1	1	1	Putative protein	GI:16411641	Strongly similar to transmembrane transporter	2×10^{-34}
At3g08040	1	0	0	0	Putative integral membrane protein		·	
Unknown At1g28070	^	0	4	4	Llum athratical mustain			
0	0 0	0 0	1 1	1 1	Hypothetical protein			
At1g28700 At1g75810	0	0	1	1	Unknown protein Hypothetical protein			
At2g40080	0	0	1	1	Unknown protein			
	0	0	1	1	Unknown protein			
At3g26000					·			
At4g32340	0	0	1	1	Unknown protein Unknown protein			
At4g12070	0	1	0	1	•			
At1g67210	0	1	1	0	Unknown protein			
At1g73330	0	1	1	1	Putative protein			
At3g47833	0	1	1	1	Unknown protein			
At3g51250	0	1	1	1	Unknown protein			
At4g27080	1	0	0	0	Unknown protein			
At4g14260	1	0	1	1	Hypothetical protein			
At3g01860	1	1	0	1	Unknown protein			

The description of the genes is retrieved from the Munich Information Center for Protein Sequences (MIPS); when the gene was annotated as unknown, the closest homolog is given as obtained from PEDANT at MIPS (Frishman et al., 2003). Expression was scored in the wild type and pal1, pal2 and pal1 pal2 double mutants (DM) with 0 for absence/decrease and 1 for presence/increase in expression, taking the wild type as reference. AGI, Arabidopsis Genome Initiative.

Table 2. Identities and Expression Patterns of the Sequence Tags Identified for Their Differential Expression in the Wild Type and *pal1* and *pal2* Mutants by Hybridization with a 6K Arabidopsis Microarray

	Expressi	on in <i>pal1</i>	Expression in pal2			Nearest Homo		
AGI	Fold Change	P Value	Fold Change	P Value	Description (MIPS)	AGI/GI	Description	E Value
Amino Acid	Polatod						·	
At3q15450	riciated		3.64	2.21 × 10 ⁻⁸	Unknown protein	At5g10240	Asn synthetase ASN3	3×10^{-7}
At4g27450			2.33	7.87×10^{-7}	Unknown protein	At5g10240	Asn synthetase ASN3	2×10^{-5}
At4g28410	0.40	4.04×10^{-8}	2.00	7.07 × 10	Hypothetical protein	At4g28420	Tyr transaminase-like	1×10^{-4}
114920410	0.40	4.04 / 10			Typothetical protein	A14920420	protein	1 × 10
At3g25990	0.46	4.47×10^{-7}			Putative DNA binding		protein	
					protein, GT-1			
arbohydra	te-Related							
t5g49360	2.19	1.91×10^{-4}	3.23	6.30×10^{-6}	Xylosidase			
t5g38410	2.01	9.54×10^{-8}	2.05	7.55×10^{-8}	Rubisco small chain 3b			
-					precursor			
At1g70290			2.79	2.29×10^{-6}	Putative trehalose-6-			
					phosphate synthase			
At4g36670			2.40	1.41×10^{-5}	Sugar transporter-like			
•					protein			
At3g01500			2.16	1.19×10^{-4}	Carbonic anhydrase,			
					chloroplast precursor			
At1g79550			2.07	1.02×10^{-8}	Phosphoglycerate kinase			
					(EC 2.7.2.3) like protein			
At2g39730	2.03	5.47×10^{-7}			Rubisco activase			
t3g23490	0.49	$2.40 imes 10^{-6}$			Cyanate lyase (CYN)			
Phenylpropa	anoid-Rela	ted						
At2g22990	2.29	9.35×10^{-5}			Putative Ser			
					carboxypeptidase I ^a			
At2g34660	0.42	1.15×10^{-5}			MRP-like ABC transporter			
At5g17220	0.41	6.20×10^{-4}			GST-like protein			
At5g42980			0.40	6.00×10^{-4}	Thioredoxin			
At2g37040	0.30	2.22×10^{-7}			Phe ammonia lyase (PAL1)			
At5g42800	0.26	4.96×10^{-5}			Dihydroflavonol 4-reductase			
xt5g13930	0.15	2.11×10^{-6}			Chalcone synthase			
ight-Relate	ea		0.00	0.05 × 40.6	DNA (dai-			
At2g41460			2.98	6.95×10^{-6}	DNA-(apurinic or apyrimi-			
			0.07	0.44 40 5	dinic site) Iyase (ARP)b	A.E. 45050	001074110 17	4 40 . 9
At1g68520			2.67	2.11×10^{-5}		At5g15850	CONSTANS-like 1	4×10^{-2}
V4400000			0.04	0.40 × 40-7	finger protein	A+0 04 470	Lhan Omentale	4 > 40-7
At1g29930			2.64		Unknown protein	At3g61470	Lhca2 protein	1×10^{-7}
At1g55670			2.09	4.15 × 10 '	Photosystem I subunit V			
					precursor (PsaG)			
Stress Resp	onse							
\t5g24490			2.57	1.08×10^{-4}	Unknown protein	GI:2636057	Stress-induced protein,	2×10^{-7}
							Bacillus subtilis	
At3g52470			2.55	3.35×10^{-5}	Unknown protein	At5g22200	NDR1/HIN1-like	6×10^{-6}
							protein	
\t2g40000			2.18	1.60×10^{-5}	Putative nematode-			
					resistance protein			
At1g75040			2.04	1.18×10^{-6}	Thaumatin-like protein	At5g38280	Receptor Ser/Thr	7×10^{-9}
							kinase PR5K	
ignal Trans	sduction/T	ranscriptional I	Regulation					
J		2.75×10^{-4}	U		Unknown protein	GI:28374149	Kelch-like protein 3,	9 × 10 ⁻⁶
t1g80440								

(Continued)

Table 2. (continued).

	Expression	Expression in pal1		on in <i>pal</i> 2		Nearest Homo		
AGI	Fold Change	P Value	Fold Change	P Value	Description (MIPS)	AGI/GI	Description	E Value
At1g13260			2.28	2.18×10^{-5}	Unknown protein	At5g20730	Putative protein	2 × 10 ⁻⁶⁶
At1g78830			2.02	2.16×10^{-4}	Unknown protein	At4g21370	Receptor kinase- like protein	1×10^{-102}
Miscellaneou	ıs							
At2g33830			2.23	2.48×10^{-6}	Putative auxin- regulated protein			
Unknown								
At5g21940			2.49	1.11×10^{-6}	Unknown protein			
At4g27380			2.47	7.59×10^{-10}	Unknown protein			
At2g15890			2.10	$4.30 imes 10^{-8}$	Unknown protein			
At1g14880			2.03	4.15×10^{-4}	Unknown protein			
At1g73330	0.39	1.91×10^{-7}			Putative protein			

Expression is given as fold change of wild-type expression. The description of the genes is retrieved from MIPS; when the gene was annotated as unknown, the closest homolog is given as obtained from PEDANT at MIPS (Frishman et al., 2003).

activase is upregulated in the pal1 mutant. Both are central parts of CO₂ fixation into organic carbon (Table 2, Figure 7). Enzymes, such as phosphoglycerate kinase and triose phosphate isomerase, that metabolize the phosphoglycerate generated by Rubisco, are also upregulated in pal2 mutants (Tables 1 and 2, Figure 7). SNF4 (Akin γ), upregulated in all pal mutants, plays important regulatory roles in carbohydrate metabolism in yeast and Arabidopsis (Table 1; Bouly et al., 1999). Trehalose 6-phosphate synthase, upregulated in pal2 mutants, generates trehalose 6phosphate that, in turn, has an important function in controlling the flux into glycolysis in yeast and controls carbohydrate use in plants (Table 2; Eastmond and Graham, 2003; Schluepmann et al., 2003). Invertase, upregulated in all mutants, and starch phosphorylase, upregulated in the pal1 mutant, participate in the generation of hexose phosphates from sucrose and starch degradation, respectively. It is tempting to think that some of these tags identify components that are involved in the generation of substrates for cell wall biosynthesis because cellulose synthase was also induced in the pal1 and pal1 pal2 double mutants (Table 1). β-Xylosidase, upregulated in both pal1 and pal2 mutants (Table 2), hydrolyzes β-D-xylose residues of (most likely) glucuronoarabinoxylans. The glucuronoarabinoxylans in turn are able to covalently bind lignins and to attach to cellulose (Hatfield et al., 1999).

Two smaller groups of tags relate to light signal transduction and stress responses, two processes in which the importance of flavonoids is well documented. Light-related transcripts are found preferentially in the *pal2* mutant. SPA1 and FAR are both specific signaling intermediates of phytochrome A signal transduction (Hoecker et al., 1999; Hudson et al., 1999). Two different FAR homologs, one upregulated and one downregulated in the *pal2* mutant, are found (Table 1). Moreover, the *pal2* mutant also

contains a CONSTANS-related transcription factor that is upregulated (Table 2). Two other light-related tags are mutS and DNA (apurinic or apyrimidinic) lyase, both involved in DNA repair that is probably increased because of the lack of UV light-protecting flavonoids (see below). ω -3 Fatty acid desaturase and allene oxide synthase participate and rate limit, respectively, the biosynthesis of jasmonate and are upregulated in all *pal* mutants (Table 1, Figure 7).

pal1 pal2 Double Mutants Lack Three Major Flavonol Glycosides

The alterations in gene transcription related to particular pathways prompted us to conduct a more in-depth biochemical characterization of the mutants. In accordance with the transcriptome data, and despite the near wild-type visible appearance of the mutants, severe modifications in amino acids, phenolics, and lignin accumulation were detected.

When the soluble phenolic compounds in the inflorescence stem of the three mutants and the wild type were compared by reverse-phase HPLC, significant changes in the profile were noticed. As shown in the maxplots presented in Figure 8, reduced PAL activity led to a decreased accumulation of several phenolics. Out of 89 peaks in the analyzed chromatograms, eight peaks differed significantly in abundance between the wild type and pal mutants. As shown in Table 3, total phenolics were significantly diminished to approximately one-third of the wild-type amount in the pal1 pal2 double mutants, slightly reduced to 75% in pal1 mutants, and not affected in the pal2 mutant. This decrease in total phenolics correlates with the levels of PAL activity (Figure 4) and is mainly attributable to the depletion of the three major peaks denoted as A, B, and C in Figure 8. As shown in Figure 9A, these

^a Note that this gene is annotated as Ser carboxypeptidase at MIPS but has been shown to correspond to the *SNG1* locus, encoding sinapoylglucose:malate sinapoyltransferase (Lehfeldt et al., 2000).

b It cannot be ruled out that the upregulation of ARP partly results from fragments of the ARP gene used in the promoter-trap construct.

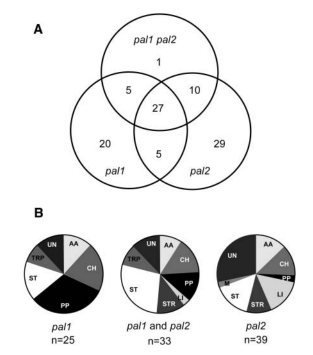


Figure 6. Number and Functional Categories of Genes with Altered Expression in *pal1*, *pal2*, and *pal1* pal2 Mutants as Compared with the Wild Type.

(A) Summary from both cDNA-AFLP and microarray analyses (Tables 1 and 2) corrected for the genes that were revealed by both methods. (B) Functional categories of the groups of genes controlled either by pal1, pal2, or both. The group pal1 comprises genes differentially expressed in only pal1 (n=20) or pal1 and the double mutant (n=5). The group pal2 comprises genes differentially expressed in only pal2 (n=29) or pal2 and the double mutant (n=10). The group pal1 and pal2 comprises genes differentially expressed in both single mutants but not in the double mutant (n=5), genes expressed in pal1, pal2, and the double mutant (n=27), and genes expressed exclusively in the double mutant (n=1). AA, amino acid metabolism; CH, carbohydrate metabolism; LI, light related; M, miscellaneous; PP, phenylpropanoid related; ST, signal transduction; STR, stress response; TRP, transport; UN, unknown.

peaks correspond, on the basis of their retention time (RT), photodiode array, and mass spectrometry (MS) spectra, to previously identified kaempferol glycosides (Veit and Pauli, 1999): kaempferol 3-O-β-[β-D-glucopyranosyl(1-6)-D-glucopyranoside]-7-O-α-L-rhamnopyranoside (Glc-Glc-Rha-kaempferol), kaempferol 3-O- β -D-glucopyranoside-7-O- α -L-rhamnopyranoside (Glc-Rha-kaempferol), and kaempferol 3-O- α -L-rhamnopyranoside-7-O- α -L-rhamnopyranoside (Rha-Rhakaempferol), respectively. The alterations in the accumulation of these peaks in the single mutants were not statistically significant but showed a trend to lower levels in the pal1 mutant (Table 3). These glycosides belong to the flavonol glycosides that are the major UV light-absorbing metabolites in Arabidopsis (Graham, 1998). Interestingly, other UV light-protecting compounds in Arabidopsis, such as sinapoyl glucose and sinapoyl malate (Landry et al., 1995), were not altered in leaves or inflorescence stems (thin-layer chromatography; data not shown). To address whether the reduction of the three major peaks in the double mutant was a result of a reduced synthesis of *trans*-cinnamic acid, double mutants and the wild type were fed for 24 h with *trans*-cinnamic acid or *para*-coumaric acid. Kaempferol glycoside peaks could not be restored in the double mutant (data not shown), suggesting that the reduced PAL activity has not only simple effects on the amount of its reaction product but imbalances the phenylpropanoid pathway in a more sophisticated way.

Feruloyl Malate Esters Coupled to Coniferyl Alcohol Are Less Abundant in pal1 and pal1 pal2 Double Mutants

Besides the three major UV light-absorbing kaempferol glycosides, five other peaks were significantly altered in the mutants, one of which occurred only in the double mutant (RT 8.350 min; not identified, Table 3). The compound at RT 6.033 min (Figure 8) was identified as glycosylated vanillic acid by RT, photodiode array spectrum, and comparison with an authentic standard (Table 3).

The remaining three peaks were significantly reduced in *pal1* and the double mutant, but not in the *pal2* mutant (RT 7.331 min, 11.178 min, and 15.119 min; Table 3). MS analysis of the compound eluting at 7.331 min indicated that this compound is scopolin or isoscopolin (Figure 9B). The peak was identified as scopolin because its aglycon released after hydrolysis consisted exclusively of scopoletin (RT 11.37 min) and not of isoscopoletin (RT 10.78 min). Scopolin had been detected previously in tobacco leaf extracts (Baumert et al., 2001) but not in Arabidopsis.

The compound eluting at 11.178 min has a UV/VIS spectrum characterized by absorption maxima at 235 and 329 nm. MS analysis suggested that this peak corresponds to the 4-O-8cross-coupling product of feruloyl malate and coniferyl alcohol, **FM**(4–O–8)**G**, or 3-{4-[2-hydroxy-2-(4-hydroxy-3-methoxy-phenyl)-1-hydroxymethyl-ethoxy]-3-methoxy-phenyl}-acrylic acid (Figure 9C). The core structure **FA**(4-O-8)**G** (FA, ferulic acid) was authenticated using previously isolated FA(4-O-8)G from ryegrass (Lolium perenne) (Ralph et al., 1992). The UV/VIS spectrum (λ_{max} at 242 and 332 nm) of the third compound (RT 15.119 min) was similar to that of FM(4-O-8)G. Based on MS analysis and its partial authentication with FA(5-8)G, this compound is suggested to be the related FM(5-8)G compound 2-{3-[2-(4-hydroxy-3-methoxy-phenyl)-3-hydroxymethyl-7-methoxy-2,3-dihydro-benzofuran-5-yl]-acryloyloxy}-succinic acid (Figure 9D). Thus, the two latter products resulted from coupling of coniferyl alcohol (at its favored 8-position) with feruloyl malate (at its 4-O- or 5-position) and had not been described before.

In summary, only downregulation of both PAL1 and PAL2 leads to significant changes in the accumulation of the major soluble phenolic compounds, the kaempferol glycosides (Table 3). On the other hand, the reduced accumulation of compounds produced by other side branches of phenylpropanoid metabolism, such as scopolin (hydroxycoumarin), and the feruloyl malates coupled to coniferyl alcohol were clearly linked with the mutation in *PAL1* (Table 3, Figure 9). Sinapate esters were

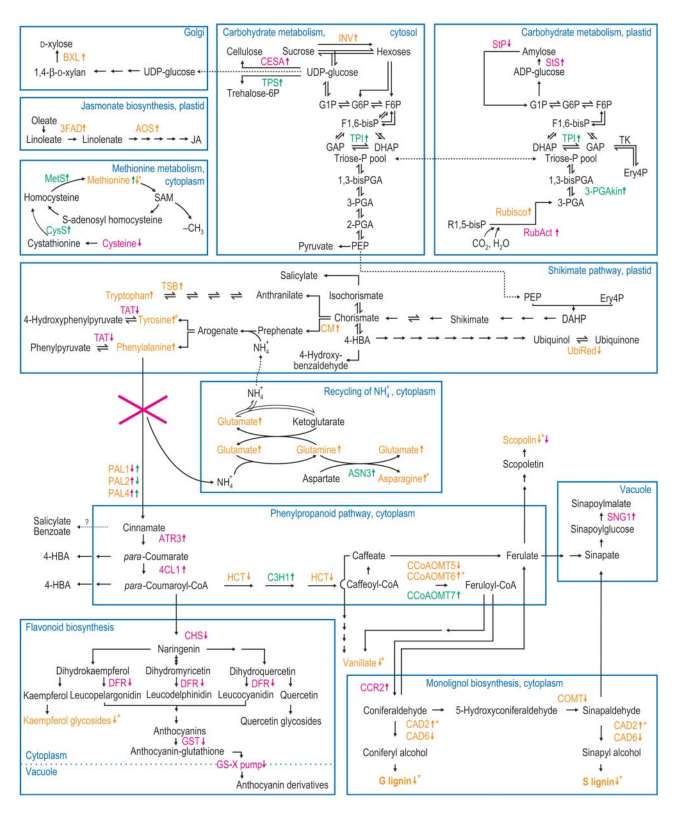


Figure 7. Integration of the Metabolic Pathways Affected by a Decreased Carbon Flux into the Phenylpropanoid Pathway, as Evidenced by Molecular Alterations in pal Mutants.

notably unaffected in all mutants, indicating that either this pathway is essential or that the enzymes producing sinapate esters are in metabolic complexes with PAL isoforms other than PAL1 and PAL2. This observation suggests that PAL1 and PAL2 are either associated with different cell types or with different metabolic complexes within the cell or that PAL1 has a higher activity than PAL2.

pal1 pal2 Mutants Overaccumulate Phe and Are Perturbed in Other Nonaromatic Amino Acids

Given the high number of differential transcripts related to amino acid metabolism, amino acid concentrations were determined in the wild type and the *pal* mutants. As shown in Table 4, the reduction in PAL activity had dramatic consequences for the accumulation of several amino acids, not only the aromatic ones. The content of most amino acids was increased in all mutants. As expected, Phe accumulated to >100-fold in the double mutant as compared with the wild type, whereas the other two aromatic amino acids, Tyr and Trp, accumulated to fourfold (Table 4). Accordingly, the *CM* gene, the first committed enzyme toward Phe and Tyr, as well as the *TSB* gene, were only expressed in the three mutants but were undetectable in the wild type in the transcript profiling experiment (Table 1).

Another affected amino acid was Met that was significantly increased in the *pal2* mutant but decreased to 6% of the wild-type level in the double mutant (Table 4). A transcript corresponding to Met synthase was exclusively upregulated in the *pal2* mutant (Table 1). Finally, and noteworthy, Gly was only detected in the wild type (Table 4). Four transcripts related to Gly degradation were exclusively expressed or upregulated in the *pal* mutants (Tables 1 and 2). By contrast, Asn was only detected in the double mutant (Table 4). Correspondingly transcripts encoding putative proteins with strong similarity to ASN were upregulated in *pal2* and *pal1 pal2* mutants (Tables 1 and 2).

In conclusion, blocking PAL led to a strong overaccumulation of Phe and imbalanced the abundance of many other amino acids. Quite surprisingly, 15 out of 20 amino acids changed significantly in at least one genotype (Table 4). The fact that the concentration of these amino acids increases, except for Met in the double mutant and Cys in *pal1*, suggests an overarching mechanism for homeostasis of amino acid levels that acts across individual biosynthetic pathways. Similarly, plants overexpressing a Trp or Tyr decarboxylase, thus artificially creating a metabolic sink, were altered in the aromatic and three nonaromatic amino acids (Yao et al., 1995; Guillet et al., 2000).

pal1 pal2 Mutants Have Less Lignin with a Higher Syringyl/Guaiacyl Ratio

To determine lignification changes, lignin contents were measured via the acetyl bromide method, and monomer compositions of the wild type and pal mutants were determined by derivatization followed by reductive cleavage (DFRC) analysis (see Methods). As presented in Table 5, inflorescence stems of 3-month-old wild-type and mutant plants clearly differed in lignin content as well as monomer composition despite the fact that these differences were hardly apparent in histochemical analysis (Figure 1). Double mutants had 30% residual lignin (Table 5), and the molar ratio of S units in the noncondensed lignin fraction was nearly double that in the wild type. S-rich lignins have higher cleavable β-aryl ether levels and less condensed structures, involving 5-5-, 8-5-, and 4-O-5-units (Ralph et al., 2004), explaining the higher amount of G and S monomers from the noncondensed fraction per gram of lignin. A similar reduction in total lignin and increase in the S/G ratio was found in transgenic tobacco downregulated for PAL (Sewalt et al., 1997; Korth et al.,

The determination of lignin content by acetyl bromide leads to an underestimation of the amount of monomers linked by ester bonds. Therefore, isolated cell wall material was saponified and analyzed by HPLC. The HPLC profiles of the wild type and double mutant were virtually identical (data not shown), indicating that the amount of esters incorporated into the cell wall was not significantly changed.

Figure 7. (continued).

Results of the transcriptome analyses (RT-PCR, cDNA-AFLP, and microarray) and the biochemical analyses (extractable phenolics, amino acids, and lignin) are incorporated. A dashed line indicates transport into another compartment; right and left harpoons mark a reversible enzyme reaction and arrows an irreversible enzyme reaction, respectively; question mark denotes a not fully established path. Arrows pointed up or down indicate the upregulation or downregulation of the gene or metabolite, respectively. The following color code is used: red, transcripts and metabolites linked with the pal1 but not the pal2 mutation; green, transcripts and metabolites linked with the pal2 but not the pal1 mutation; orange, transcripts and metabolites linked with the pal1 and pal2 mutations; orange with asterisk, transcripts and metabolites occurring only in the pal1 pal2 double mutant. Abbreviations are listed per pathway. Carbohydrate metabolism: 1,3-BisPGA, 1,3-bisphosphoglycerate; BXL, β-xylosidase; CESA, cellulose synthase; DHAP, dihydroxyacetone phosphate; Ery4P, erythrose 4-phosphate; F1,6-BisP, fructose 1,6-bisphosphate; F6P, fructose 6-phosphate; G1P, glucose 1-phosphate; G6P, glucose 6-phosphate; GAP, glyceraldehyde 3-phosphate; INV, invertase; PEP, phospho-enol-pyruvate; 2-PGA, 2-phosphoglycerate; 3-PGA, 3-phosphoglycerate; 3-PGAkin, 3-phosphoglycerate kinase; R1,5-BisP, ribulose 1,5-bisphosphate; RubAct, Rubisco activase; StP, starch phosphorylase; StS, starch synthase; TK, transketolase; TPI, triose-phosphate isomerase; TPS, trehalose 6-phosphate synthase. Jasmonate biosynthesis: AOS, allene oxide synthase; 3FAD, ω-3 fatty acid desaturase; JA, jasmonate. Methionine metabolism: CysS, cystathionine β-lyase (Cys synthase); MetS, Met synthase. Shikimate pathway: DAHP, 3-deoxy-D-arabino-heptulosonate 7-phosphate; Ery4P, erythrose 4-phosphate; 4-HBA, para-hydroxybenzoate; 4-HBald, para-hydroxybenzaldehyde; PEP, phospho-enol-pyruvate; UbiRed, NADH-ubiquinone oxidoreductase. Recycling of NH₄+1: ASN3, Asn synthase 3. Phenylpropanoid pathway: ATR3, NADPH-ferrihemoprotein reductase 3; CCoAOMT, caffeoyl-CoA 3-O-methyltransferase; 4-HBA, para-hydroxybenzoate. Monolignol biosynthesis: G lignin, guaiacyl lignin monomers; S lignin, syringyl lignin monomers. Flavonoid biosynthesis: CHS, chalcone synthase.

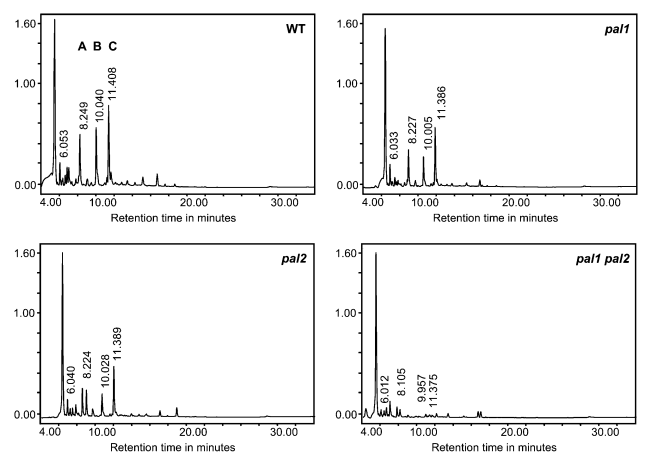


Figure 8. Profiles of Soluble Phenolic Metabolites in 3-Month-Old Inflorescence Stems of the Wild-Type C24, *pal1*, and *pal2*, *pal1 pal2* Mutants. Soluble phenolics are separated by reverse-phase HPLC. The maxplot chromatograms are given as absorbance units at 200 to 450 nm and are scaled down to show the differences in the most abundant peaks. A, B, and C denote the three major kaempferol glycosides decreased in the double mutant.

DISCUSSION

Molecular Phenotyping Is Informative

Genome-wide studies of the transcriptome allow explaining a mutant phenotype at the molecular level at a depth that was not possible before the development of genomic tools. The extension of the phenotypic description to the molecular level increases the resolution so that functional redundancy of closely related gene family members can be separated. Thus, these tools are excellent to provide an insight into the functional significance of individual gene family members. Furthermore, our study underscores that the disruption of a single gene can have wide-ranging consequences on the transcription of other genes, revealing the intricacy of interactions among metabolic pathways, even in the absence of a visible phenotype (Tables 1 and 2). Elements of most of the pathways, altered in pal mutants, have been previously described for their responsiveness to pathogen attack or elicitation (Figure 7; Somssich and Hahlbrock, 1998). However, in contrast with transcriptome profiling after challenging the plant, typically designed to reconstruct a pathway response to a particular agent or environmental stress,

our aim was to describe the adaptation of the transcriptome to genetic differences at a particular locus. In a similar study on the brevipedicellus mutant that is disrupted in a KNOX gene and displays premature lignin deposition, expression differed in 12 genes related to lignification or cell wall biosynthesis (Mele et al., 2003). Among these are 4CL1, PAL1, TAT, and cellulose synthase, genes that are also differentially expressed in the pal mutants (Tables 1 and 2). Moreover, ASN, differentially expressed in the brevipedicellus and pal2 mutant, is known to respond dramatically at the transcriptional level to metabolites, light, and development (Lam et al., 1998). Thus, our study might have identified genes that are under metabolic regulation, as opposed to genes involved in a pathway's response to environmental or developmental signals.

Altogether, the *pal1* and *pal2* mutations cause 58 and 72 genes, respectively, to change in expression within our experiment and degree of transcriptome resolution (20% with the microarray and 60% with the cDNA-AFLP; Figure 6). In the *pal* mutants, the expression of several differentially expressed genes depends probably on the level of one or more biochemical compounds generated or consumed by PAL, such as Phe and cinnamic acid or their upstream and downstream derivatives.

		Peak Height in $\mu V/\mu g$ Dry Weight (Average with SE)					
RT (min)a	Identity	Wild Type	pal1	pal2	pal1 pal2		
Total		99.87 ± 6.65	73.98 ± 8.43	106.01 ± 12.95	34.14 ± 5.73*		
6.033	Glycosylated vanillic acid	5.89 ± 0.29	5.65 ± 0.19	6.08 ± 0.99	$2.26 \pm 0.34^*$		
7.331	Scopolin	0.67 ± 0.07	$0.41 \pm 0.02^*$	0.64 ± 0.09	$0.19 \pm 0.04^*$		
8.227	Glc-Glc-Rha-kaempferol	19.12 ± 2.07	12.95 ± 2.21	17.49 ± 2.62	0.21 ± 0.08*		
8.350	Not identified	ND	ND	ND	0.03 ± 0.01		
10.005	Glc-Rha-kaempferol	20.67 ± 2.59	12.61 ± 3.35	19.39 ± 3.31	$0.31 \pm 0.10^*$		
11.178	FM (4-O-8) G ^b	1.21 ± 0.15	$0.62 \pm 0.06^*$	0.83 ± 0.11	$0.22 \pm 0.05^*$		
11.386	Rha-Rha-kaempferol	25.89 ± 2.32	19.71 ± 3.02	27.66 ± 3.70	$0.64 \pm 0.23^{*}$		
15.119	FM (5-8) G °	1.34 ± 0.13	$0.74 \pm 0.09^*$	1.16 ± 1.27	$0.42 \pm 0.09^*$		

The $10^6 \mu V$ corresponds to an absorbance of 1 OD. One-way ANOVA and post-hoc least square difference tests for peak height were performed (see Methods) and are marked by an asterisk, if found significantly different from the wild type at P = 0.05. ND, not detected.

However, among the differentially expressed genes, there will also be genes that are expressed as a result of secondary effects of altered cellular processes.

A Reduced Carbon Flux into the Phenylpropanoid Pathway Affects the Homeostasis of Amino Acid and Carbohydrate Metabolism

Under normal conditions, 20% of all fixed carbon is pumped into the shikimate pathway (Herrmann, 1995). One of the major drains to this pathway is PAL, consuming Phe. Because the *pal1 pal2* double mutant has 30% residual total extractable phenolics and 30% residual lignin, carbon supply must be either adjusted or diverted to other pathways (Tables 3 and 5). Taking into account, moreover, that *PAL1* expression occurs only in the vascular bundle (Leyva et al., 1995) and that *PAL2* expression also preferentially correlates with lignified inflorescence stems (Raes et al., 2003), most of the observed molecular alterations in the mutants are expected to be manifest in a limited number of cell types in the stem (Figure 1).

Assuming that most transcriptomic changes concern the same few cell types, the three major downstream branches of the phenylpropanoid pathway (flavonoids, lignins, and sinapate esters) undergo a different adaptation to the reduced flux into the pathway in the mutants: the production of flavonoids and lignin is clearly reduced. Additionally, hydroxycoumarins, glycosylated vanillic acid, and **FM**(4–O–8)**G** and **FM**(5–8)**G** are reduced in abundance (Table 3), whereas sinapate esters remain unchanged. Because all flavonoids, sinapate esters, and hydroxycoumarins have UV light-protecting capacities, the sinapate esters remain probably produced because they are most effective in UV light shielding. Alternatively, sinapate esters could be derived from a metabolic complex involving a PAL isoform other than PAL1 or PAL2.

As summarized in Figure 7, signs of molecular adjustments are not only found in the phenylpropanoid pathway but also in the shikimate pathway, in the related cell wall biosynthesis, in components of central metabolism such as glycolysis, and even in distant stress-related pathways. In the transcriptome data (Figures 3 and 5, Tables 1 and 2), the complete sequence of the early phenylpropanoid biosynthesis is reconstructed, except for C4H (Figure 7). Similarly, in PAL-suppressed tobacco plants, *C4H* expression remains unchanged (Blount et al., 2000). However, a NADPH P450 reductase gene *ATR3*, shown to support C4H activity in vitro, was upregulated (Table 1; Urban et al., 1997; Mizutani and Ohta, 1998; Ro et al., 2002). Probably the reduced flux through PAL superinduces transcription of the consecutive genes: the respective other *PAL* gene(s), *ATR3*, *4CL1*, and *C3H1*. To the contrary, genes encoding components in the flavonoid pathway (e.g., *CHS*, *DFR*, *GST*, and *AtMRP2*) and genes thought to be specific for lignification (e.g., *COMT* and *CAD6*) are downregulated.

With respect to the shikimate pathway generating the aromatic amino acids, transcripts for the enzyme at the branch point (CM) and two enzymes at the respective end points (TSB and TAT) were differentially expressed (Figure 7). All three hold a great potential for exerting control over the flux in this pathway. Phe, accumulating to 100-fold in the double mutant, and Trp, accumulating to fourfold in the double mutant, exert negative and positive feedback control on CM, respectively, and Trp also negatively regulates anthranilate synthase in its own biosynthesis (Bentley, 1990). On the other hand, Trp synthase is upregulated in pal mutants, pointing to a counterbalancing mechanism for decreased anthranilate synthase. TAT, transcriptionally downregulated in pal1 mutants, degrades Phe and Tyr (Figure 7). Its expression might be downregulated to limit the degradation of Phe to keep feeding substrate into the PAL reaction. Together, a severe imbalance in a part of the pathway might override classical feedback mechanisms. The accumulation of aromatic amino acids seems to cause further adjustments in the abundance of as many as 15 amino acids, indicating a general amino acid homeostasis (Table 4).

Clearly, many changes also occur in the primary metabolism of the *pal* mutants (Figure 7). Several studies reveal the reverse,

^a The RTs correspond to those in an individual pal1 chromatogram, shown in Figure 8.

^b Feruloyl malate coupled 4–O–8 to coniferyl alcohol.

^c Feruloyl malate coupled 5-8 to coniferyl alcohol.

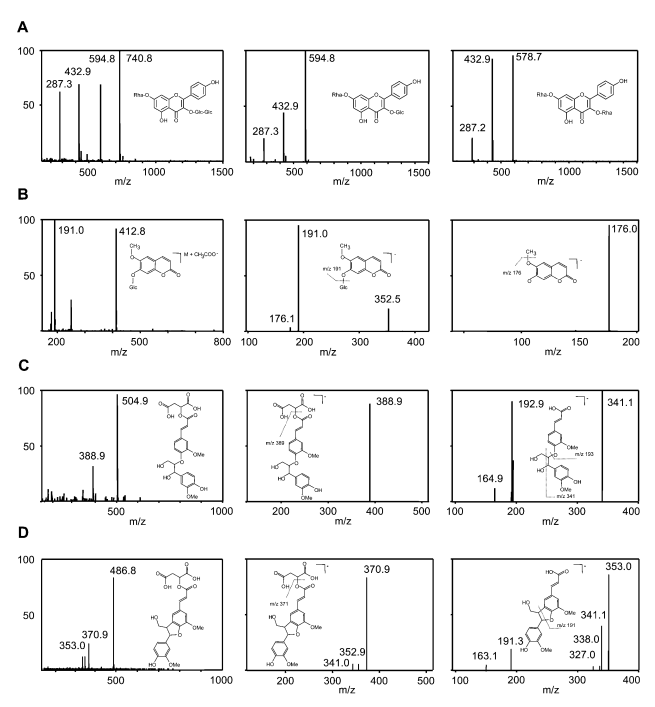


Figure 9. Identification by Mass Spectrometry of Five Soluble Phenolics Significantly Altered in pal Mutants.

All MS spectra are given as relative abundance of molecules with a scale to 100%.

(A) Full MS spectra obtained in the positive ionization mode for the three major peaks found to be altered significantly in *pal* mutants (indicated with A, B, and C in Figure 8). The mass-to-charge (m/z) values in the MS spectra for A, B, and C confirm that these compounds are hexose-conjugated kaempferols. The A in Figure 8 corresponds to kaempferol $3-O-\beta-[\beta-D-glucopyranosyl (1-6)-D-glucopyranoside]-7-<math>O-\alpha-L$ -rhamnopyranoside, B to kaempferol $3-O-\beta-D-glucopyranoside-7-O-\alpha-L$ -rhamnopyranoside, and C to kaempferol $3-O-\alpha-L$ -rhamnopyranoside-7- $O-\alpha-L$ -rhamnopyranoside.

(B) Identification of the HPLC peak at RT 7.331 min as scopolin by full MS in the negative ionization mode, MS² of m/z 413, and MS³ of m/z 191.

(C) Identification of the HPLC peak at RT 11.178 min as feruloyl malate coupled 4–O–8 to coniferyl alcohol by full MS, MS² of m/z 505, and MS³ of m/z 389. The MS⁴ of m/z 193, with the characteristic fragmentation pattern of ferulic acid, is not shown.

(**D**) Identification of the HPLC peak at RT 15.119 min as feruloyl malate coupled 5–8 to coniferyl alcohol by full MS, MS² of m/z 487, and MS³ of m/z 371. Fragmentation of m/z 371 yielded peaks at m/z 353, 341, and 338, in abundances reminiscent of a phenylcoumaran (K. Morreel and W. Boerjan, unpublished data).

Table 4. Concentrations of Amino Acids in 3-Month-Old Inflorescence Stems of the Wild Type and pal1, pal2, and pal1 pal2 Mutants as Determined by HPLC

		Peak Height in μAU/	μg Dry Weight (Average	with SE)	
Amino Acid	RT (min)	Wild Type	pal1	pal2	pal1 pal2
Asp	3.75	567.0 ± 65.6	741.3 ± 35.2	722.7 ± 38.7	930.7 ± 182.6
Glu	4.00	1185.8 ± 133.8	1744.0 ± 42.1	1745.2 ±83.3	2728.5 ± 527.3*
Asn	6.05	ND	ND	ND	4.2 ± 4.2
Ser	6.20	117.0 ± 24.7	147.7 ± 5.7	184.3 ± 11.4	177.6 ± 25.8
Gln	6.48	78.3 ± 16.1	135.8 ± 4.6	211.2 ± 29.8	741.2 ± 149.0*
Gly	6.65	1.1 ± 1.1	ND	ND	ND
His	7.20	8.3 ± 2.0	15.1 ± 3.4	15.9 ± 1.1	25.5 ± 6.1*
Arg	8.10	ND	ND	ND	ND
Thr	8.30	101.7 ± 15.3	200.2 ± 5.5	181.8 ± 12.4	536.7 ± 98.3*
Ala	8.65	97.0 ± 21.2	128.1 ± 2.6	191.9 ± 14.0	$300.8 \pm 40.1^*$
Pro	8.95	23.3 ± 4.7	58.0 ± 3.3	47.0 ± 6.1	198.2 ± 58.8*
Tyr	11.20	45.0 ± 13.5	58.4 ± 4.6	50.1 ± 6.1	166.0 ± 31.2*
Val	11.85	112.1 ± 18.3	191.3 ± 5.0	237.1 ± 7.6	494.6 ± 78.3*
Met	12.20	439.1 ± 41.8	522.7 ± 34.9	717.1 ± 117.2*	$26.3 \pm 3.2^*$
Cys	12.70	7.2 ± 1.2	1.1 ± 1.1*	6.8 ± 0.7	8.2 ± 0.8
lle	13.45	40.1 ± 7.2	77.1 ± 0.9	86.4 ± 6.7	161.9 ± 23.7*
Leu	13.65	43.6 ± 5.9	91.8 ± 4.7*	70.3 ± 3.8	103.9 ± 14.9*
Phe	14.35	19.9 ± 4.1	72.8 ± 2.4	60.9 ± 4.9	2653.3 ± 740.7*
Trp	14.62	27.3 ± 3.1	49.2 ± 3.3	67.4 ± 5.6	121.3 ± 31.1*
Lys	15.30	11.8 ± 3.6	$24.7 \pm 0.8^*$	18.2 ± 1.6	21.9 ± 2.6*

Data are average of five individuals per genotype. One-way ANOVA and post-hoc Dunnett's tests were conducted (see Methods). Values significantly different from the wild type at P < 0.05 are indicated by asterisks. ND, not detected; AU, absorption units.

namely, consequences of downregulation or disruption of enzyme functions in primary metabolism for secondary metabolism. For example, downregulation of 3-deoxy-D-arabino-heptulosonate 7-phosphate synthase (shikimate pathway) or plastid transketolase (TK, Figure 7) leads to a decrease in aromatic amino acids, phenylpropanoids, and lignin (Jones et al., 1995; Henkes et al., 2001). On the contrary, here, far-reaching consequences for primary metabolism caused by a disruption in an enzyme of secondary metabolism have been found at the transcriptional level (Figure 7). One striking example is the upregulation of the Rubisco-encoding gene in pal1 and pal2 mutants (Table 2). Only the opposite (namely, Rubisco downregulation and a corresponding decrease in the Phe pool) have been shown before in tobacco (Matt et al., 2002). These observations raise the possibility that primary and secondary metabolism are interlocked through Phe pools.

Reduction in Lignin Is Not Compensated for by Cellulose in pal Mutants

Lignin and polysaccharides are two principal components of the secondary thickened cell wall. A common assumption is that these cell wall components are coordinately regulated and can compensate for each other. In the pal mutants, having reduced lignin levels, several transcripts related to cell wall polysaccharides (most importantly, cellulose synthase and β -xylosidase) were identified as upregulated, but the corresponding changes in cell wall composition could not be detected. No clear differences between the wild type and mutants were found by cellulose histochemistry and analysis of the total polysaccharide compo-

sition (data not shown). Thus, the decrease in lignin does not seem to be compensated for by cellulose. Arabidopsis mutants with a reduced cellulose synthesis can exhibit ectopic lignification (Taylor et al., 1999; Caño-Delgado et al., 2000, 2003), whereas irregular xylem 4 mutants, disrupted in the CCR gene, have less lignin but not more cellulose (Jones et al., 2001). Thus, a reduction in cellulose synthesis, which precedes lignin deposition, is sensed to induce extra lignification (Ellis et al., 2002; Caño-Delgado et al., 2003), but a reduction in lignin deposition is not necessarily compensated for by cellulose biosynthesis. In a few cases, in which a reduced lignin accumulation was accompanied by an increased cellulose content, alterations in development have been observed (Hu et al., 1999; Li et al., 2003). An increased cellulose content in lignin-deficient plants might arise from pleiotropic effects of decreased lignification rather than from compensation for lignin by cellulose in each single cell. Alternatively, the lack of polysaccharide changes in the pal mutants has to be explained by the subtle nature of these changes or the failure of the methods applied; a complex mixture of different cell types present in the whole inflorescence stem is used for chemical analyses, whereas only the fraction of cells with secondary cell wall thickening is expected to display changes.

A Role in Cross-Linking for the Novel Coupling Products of Feruloyl Malate and Coniferyl Alcohol?

Although no gross differences were observed, various signs of decreased cell wall rigidity, particularly in the interfascicular fibers, were noticed in the double mutant (Figure 2). This

Table 5. Lignin Content and Monomer Composition in Inflorescence Stems of 3-Month-Old Wild-Type, pal1, pal2, and pal1 pal2 Mutants

Genotype	mg Lignin/g Cell Wall	μmol G/g Lignin	μmol S/g Lignin	Molar Ratio S/G
Wild type	200 ± 8	761 ± 103	154 ± 9	0.21 ± 0.02
pal1	139 ± 2	715 ± 56	197 ± 15	0.28 ± 0.003
pal2	125 ± 3	755 ± 63	216 ± 11	0.30 ± 0.03
pal1 pal2	69 ± 1	1020 ± 30	389 ± 33	0.38 ± 0.04

Lignin content was determined as acetyl bromide soluble lignin (ABSL). Monomer compositions were determined by DFRC (see Methods). Data were obtained with at least three individuals/genotype; each individual was measured twice. Data are expressed as averages with SE.

phenotype is probably linked to the 30% residual lignin levels in the pal1 pal2 mutants but could possibly also arise from the decreased abundance of the novel compounds, FM(4-O-8)G and FM(5-8)G, in the pal1 and pal1 pal2 mutants (Table 3). The malate esters of FA coupled to coniferyl alcohol have not yet been identified before in plants, but ajugol, methyl, and glucose esters of FA(4-O-8)G and/or FA(5-8)G have been reported (Yamamoto et al., 1993; Wahl et al., 1995; Liu et al., 1999). The core structure FA(4-O-8)G esterified to arabinosyl units has been found in saponified grass and cereal cell walls (Jacquet et al., 1995; Grabber et al., 2002; Bunzel et al., 2004). Generally, FA is esterified to the polysaccharides during their biosynthesis in the endomembrane system, and the feruloyl esters can undergo oxidative coupling, thereby linking polysaccharides in the cell wall (Ralph et al., 1999; Fry, 2004). It is possible that the malate ester of FA (coupled with coniferyl alcohol) is an intermediate that can become transesterified to polysaccharides. If this were the case, the reduced rigidity of the cell wall observed in the double mutant could partly arise from weaker crosscoupling of polysaccharides with each other and with lignin.

The Molecular Phenotypes of pal1 and pal2 Mutants Reveal Distinct Roles for PAL1 and PAL2 in Phenylpropanoid Production

Isoforms of PAL have been shown to differ in posttranslational modification, metabolite sensitivity, localization, and association with metabolite channeling complexes (Sarma et al., 1998; Allwood et al., 1999; Rasmussen and Dixon, 1999; Kao et al., 2002). Only few studies have examined the contribution of isoforms to different branches of the phenylpropanoid pathway (Howles et al., 1996; Sarma et al., 1998; Rasmussen and Dixon, 1999). However, the specificity of the isoforms from other species cannot readily be transferred to those in Arabidopsis.

pal1 and pal2 mutants are indistinguishable at the phenotypic level, and only the double mutant has slight phenotypic changes together with a clear reduction in kaempferol glycosides and lignin (Figures 1 and 2, Tables 3 and 5). Considering the 34 genes that code for the enzymes of monolignol biosynthesis, similar trends in alteration of gene expression were observed for most of them in both single pal mutants (e.g., HCT, CCoAOMT5, COMT, and CAD6, Figure 5). Although this observation argues for a functional overlap of PAL1 and PAL2, the fact that expression levels of these four genes were closer to the wild-type level in the double mutant than in the single mutants corroborates a more complex metabolic interaction (Figure 5). On the other hand,

expression of several genes was only significantly increased in the double mutant (*4CL1* and *CCoAOMT6*; Figure 5), arguing for a synergistic action of PAL1 and PAL2 in the generation of particular phenylpropanoids. Still, the differential expression of several genes is correlated with only one of the two mutations: *PAL3*, *C3H1*, and *CCoAOMT7* with *pal2*, and *CCR2* with *pal1* (Figures 3 and 5). Although it is too early to speculate on how this transcriptional regulation corresponds to the differential accumulation of a particular metabolite or with particular metabolic complexes, these transcriptional differences clearly indicate directions toward the specific roles of these two PAL isoforms.

When the altered transcripts of other pathways in either *pal1* or *pal2* (and the double mutant) are compared, more tags relate to light, UV light protection, and stress in *pal2* than in *pal1* mutants (Figures 5, 6, and 7). On the other hand, *pal1* mutants are more altered in phenylpropanoid-related transcripts (*PAL1*, *PAL2*, *ATR3*, *4CL1*, *DFR*, *CHS*, *GST*, *GS-X pump*, and *SNG1*, Figure 7). In accordance with these transcriptomic changes, the *pal1*, but not the *pal2*, mutant has reduced total extractable phenolics because of a decrease in sugar-conjugated kaempferols, scopolin, **FM**(4–O–8)**G**, and **FM**(5–8)**G** (Table 3).

The difference in aromatic carbon flux seems to translate in different ways to carbon flux in both mutants. The lack of C6-C3 carbon probably signals in both mutants a demand for more carbon; transcripts correlated with photosynthetic carbon fixation and with sucrose degradation are upregulated in both mutants (e.g., Rubisco and invertase; Figure 7). However, based on the transcriptomic changes that exclusively occur in pal1 mutants, the excess carbon is stored away as starch in pal1 mutants (starch synthase: up, starch phosphorylase; involved in starch degradation: down; Figure 7). On the other hand, in pal2 mutants, carbon is rather redirected toward other processes; transcripts related to glycolysis and gluconeogenesis (e.g., trehalose 6P-synthase, triose-P isomerase, phosphoglycerate kinase: up; Figure 7). Possibly part of this carbon is used to generate more amino acids because the levels of many amino acids are higher in pal2 than in pal1 (Table 4).

The understanding of the transcriptome and biochemistry provide novel insight into the functional differences of the two PAL isoforms. On the basis of these observations, a PAL1 knock down has more severe consequences for the phenylpropanoid pathway than a PAL2 knock down. Whereas both enzymes contribute to the production of lignin precursors, PAL1 is of higher importance for the generation of flavonoids in the inflorescence stem. To elaborate on the specific functions of the PAL isoforms in Arabidopsis, future experiments need to address

the function of the two other PALs and the potentially different localization of all PALs at the cellular and tissue levels.

The application of a similar molecular phenotyping strategy to other mutants in the 34 genes that encode the 10 known enzymes in the monolignol pathway (Raes et al., 2003) will allow us to answer long-standing questions on the metabolic regulation within the phenylpropanoid pathway, between metabolic pathways, and on the functional divergence in the gene families. In addition, comparative transcriptome analysis of this set of mutants should allow the discrimination between genes regulated by metabolites and genes expressed as a result of pleiotropic effects of a particular mutation. The comparative approach will also help to elucidate the function of the class of unknown genes that still constitute 20% of all differential transcripts identified in the pal mutants. These unknown genes are expected to fall into the same broad functional categories as the other differential genes; hence, new candidate genes that play a role in phenylpropanoid biosynthesis will be identified.

METHODS

Plant Growth Conditions

For germination, seeds were surface sterilized and placed on MS medium (Duchefa, Haarlem, The Netherlands) supplemented with 10 g L $^{-1}$ sucrose (and if required with 50 mg L $^{-1}$ kanamycin). After the seeds had undergone a cold treatment for homogenous germination (overnight at 4°C), they were exposed to 20°C , 50 μ mol m $^{-2}$ s $^{-1}$ light intensity, and 70% humidity under a 16-h-light/8-h-dark cycle. Plants were transferred to a greenhouse after 14 d. Conditions were as follows: 50 μ mol m $^{-2}$ s $^{-1}$ light intensity at plant level (MBFR/U 400-W incandescent lamps; Philips, Eindhoven, The Netherlands), a 16-h-light/8-h-dark cycle, 40% relative humidity, 23°C, and without shielding from incident daylight. Plant material was always harvested between 11 AM and 1 PM.

Generation of the *pal* Mutants and Double Mutant and Identification of the Genotypes

Primary transformants of SL12-17 (pal1) and of L1-138 (pal2) were obtained through transformation with the pSL1 and pSL4 constructs, respectively (Babiychuk et al., 1997). The exon-trap vector contains, starting from the right border of the T-DNA, the first intron and second exon of the apurinic endonuclease gene fused in frame to the neomycin phosphotransferase II (NPTII) gene lacking its translation initiation codon. The first intron of the apurinic endonuclease gene lacks its 5' donor splice site. Together, kanamycin resistance after integration of this T-DNA is only achieved in those plants, where the host gene provides promoter activity, an ATG codon, and a 5' donor splice site. After the intron is spliced out, a chimeric transcript of the host gene, apurinic endonuclease exon 2, and NPTII is made. Insertions in the PAL1 and PAL2 genes were revealed by 5' rapid amplification of cDNA ends using a NPTII primer. The precise insertion place for both genes was delineated to the first intron through sequencing of a genomic PCR product spanning from the first PAL1 and PAL2 exon to the 5' end of the NPTII gene (data not shown). Single locus, homozygous lines were obtained by self-fertilization and segregation analysis.

The pal1 pal2 double mutant was generated by crossing pal1 and pal2, using either as female and male parent. In the F2 progeny of both crosses, sterile double mutants were identified in the expected Mendelian ratios. Therefore, the double mutant had to be identified for each experiment in the progeny of plants homozygous for one but heterozygous for the other

mutation. To identify the wild-type and mutated *PAL1* and *PAL2* genes, the following primers were used as sense and antisense primers, respectively: for *pal1*, 5'-ATGGAGATTAACGGGGCACAC-3' and 5'-GCATCAGAGCAGCCGATTGTCTGTT-3'; for *PAL1*, 5'-ATGGAGATTAACGGGGCACAC-3' and 5'-CATGGCGGCTCTTGTGGCGG-3'; for *pal2*, 5'-ATGGATCAAATCGAAGCAATG-3' and 5'-GCATCAGAGCAGCCGATTGTCTGTT-3'; for *PAL2*, 5'-ATGGATCAAATCGAAGCAATG-3' and 5'-CATGGCGGCTCTTGTGGCGG-3'. PCR reactions (reaction buffer as supplied with the Taq polymerase, deoxynucleotide triphosphate at 200 pmol, primers at 100 ng) were performed as follows: 30 cycles at 94°C for 30 s, at 50°C for 30 s, at 72°C for 2 min, and a final extension step at 72°C for 10 min. PCR products were separated in 1% agarose gels.

DNA and RNA Extractions

For the identification of double mutants in the progeny of plants homozygous for one but heterozygous for the other mutation, DNA was extracted for use in PCR reactions according to Edwards et al. (1991). Total RNA from stem tissues was extracted with the TRIzol method (Invitrogen, Carlsbad, CA). Total RNA (5 μ g) was subsequently reverse transcribed into double-stranded cDNA.

RT-PCRs

All primers used for the amplification of monolignol biosynthesis genes were as described in Raes et al. (2003) and are available at http://www.psb.ugent.be/bioinformatics/. All these primers spanned over at least one intron, so that amplification from the contaminating genomic template could easily be recognized. The *ACT2* gene (At3g18780), to which the RT-PCRs were normalized when expressed as relative expression levels, was amplified with the following sense and antisense primers, respectively: 5'-GTTGCACCACCTGAAAGGAAGT-3' and 5'-CAATGG-GACTAAAACGCAAAAC-3' to generate a fragment of 364 bp on cDNA.

In RT-PCR experiments, reactions in 25 μ L (reaction buffer as supplied with the Taq polymerase, 50 ng of each primer) contained a modified nucleotide mix: dCTP, dTTP, and dGTP were at 200 pmol, whereas dATP was reduced to 20 pmol. To each reaction, 0.1 μ L of 33 P-labeled dATP (10 mCi mL $^{-1}$, 2500 Ci mmol $^{-1}$) was added, resulting in a hot-to-cold dATP ratio of 1:2500. Products were separated on 4.5% polyacrylamide or 1% agarose gels and visualized on dried gels through autoradiography. To assure at least semiquantitative assays, the linear range of the PCR reaction was determined for each gene by testing at least two template concentrations (1 μ L 1:10 diluted cDNA and 1 μ L undiluted cDNA) and three different PCR reaction cycles (21, 24, and 30).

Transcript Profiling and Handling of Differentially Expressed cDNA Fragments

cDNA-AFLP starting with 5 μ g of total RNA was performed as described by Breyne et al. (2002). Amplified fragments were considered interesting when their pattern of expression was the same in the two biological sets used. Fragments were cut out, reamplified, and directly sequenced. Sequences were subjected to a tBLASTX against GenBank and a BLASTN against all *Arabidopsis thaliana* accessions at EMBL. The best-hit sequence was aligned with the fragment and analyzed for the correct BstYI and Msel restriction sites in the respective part of the sequence, for the correct nucleotide extensions at the restriction sites, and for the correct size of the resulting BstYI/Msel fragment. Only when all criteria were fulfilled was the database hit considered correct.

Microarray Hybridization

The 6528 genes were spotted in duplicate on the microarray, of which 6008 were from a unigene clone collection from Incyte (Arabidopsis Gem

I; Incyte, Palo Alto, CA) and 520 positive and negative controls from the Universal Score Card spike set (Amersham Biosciences, Little Chalfont, UK); for details, see http://www.microarrays.be/service.htm/currently available arrays. Total RNA (5 μg) of each sample was reverse transcribed and amplified according to a modified protocol for in vitro transcription and subsequently fluorescently labeled with Cy5 or Cy3 (Amersham Biosciences; http://www.microarrays.be/service.htm/protocols). Hybridization and washing were performed in an automated hybridization station (Amersham Biosciences) for a 16-h cycle (http://www.microarrays.be/service.htm/protocols). The arrays were scanned at 532 and 635 nm by a Generation III scanner (Amersham Biosciences), and images were analyzed with ArrayVision (Imaging Research, St. Catharines, Ontario, Canada). Spot intensities were measured as artifact-removed total intensities (ARVol) without correction for background.

Microarray Data Analysis

Slides were normalized by plotting for each single slide $M = log_2(R/G)$ against $A = log_2\sqrt{R} \times G$ for each spot (Yang et al., 2002). Subsequent Lowess normalization (Lowess window width = 0.2) yielded adjusted log_2R and log_2G signal intensities. Normalized spots were processed by two sequential analyses of variance (ANOVAs; Wolfinger et al., 2001). Using this mixed model approach, as implemented in restricted maximum likelihood assay in Genstat (Payne and Arnold, 2002), the fixed and random effects were fitted simultaneously, and significance for the fixed genotype effects was calculated using the Wald statistics. The P value cutoff was set at 0.001. No further adjustments for multiple testing were performed.

Determination of PAL Activity

PAL activity was determined as previously described (Legrand et al., 1976). Homogenized samples were extracted with 0.1 M sodium borate, pH 8.8, and centrifuged for 10 min at 4°C and 14,000 rpm. The total protein content of the supernatant was determined according to the Bradford assay. Samples of 10 to 50 μg total protein were then allowed to digest a mixture of 100 nmol L-Phe and 0.01 nmol $^{14}\text{C-L-Phe}$ (ratio 10,000:1 cold:hot) for 3 h at 37°C. The volume was increased by the addition of water and extracted with a double volume of cyclohexane/ether (1:1 [v/v]). The organic phase was measured in a scintillation counter for the $^{14}\text{C-}trans$ -cinnamic acid. The partitioning coefficient of cinnamic acid between the water and the organic phase was determined to be $\sim 50\%$ under our experimental conditions. Using 0.2 nmol $^{14}\text{C-L-Phe}$ as a standard for the counts/minute, the pmol cinnamic acid s $^{-1}$ μg^{-1} protein was calculated.

Light Microscopy and Lignin Histochemistry

Vibroslices (50 to 100 μm thick) of agarose-embedded fresh stem material of 3-month-old plants were collected in water or 92% ethanol and immediately stained. Lignin staining after Wiesner was performed in 1% phloroglucinol in 92% ethanol for 2 min. The specimens were subsequently incubated in 25% HCl and immediately observed. Specimens for lignin staining after Mäule were stained in 1% KMnO₄ for 5 min, rinsed with water, incubated in 37% HCl:H₂O (1:1 [v/v]) for 2 min, and observed after the addition of a drop of NH₃. Lignin staining with concentrated aniline sulfate in water proceeded for 3 to 5 min and was stopped by the addition of diluted H₂SO₄:H₂O (1:2 [v/v]). Specimens were observed in water (Spearing, 1953).

Transmission Electron Microscopy

Basal parts of 3-month-old inflorescence stems of all genotypes were harvested and trimmed to smaller pieces before fixation. These pieces were fixed with a mixture of 4% paraformaldehyde and 3% glutaralde-

hyde and postfixed with 1% OsO_4 and 1.5% $K_3Fe(CN)_6$ in 0.1 M Nacacodylate buffer, pH 7.2. Samples were dehydrated through a graded ethanol series, including a bulk staining with 2% uranyl acetate at the 50% ethanol step. Ethanol was subsequently replaced by LR white hard grade embedding medium (London Resin, Basingstoke, UK) and samples embedded in LR white. Ultrathin sections of gold interference color were cut using an ultramicrotome and collected on collodion-coated Cu grids of 200 mesh. The sections were post-stained in an ultrostainer (Leica, Herburgg, Switzerland) for 15 min in 2% uranyl acetate (Ultrostain 1; Leica) at 40°C and 4 min in lead citrate (Ultrostain 2; Leica) at 20°C. The sections were examined with a transmission electron microscope 1010 (JEOL, Tokyo, Japan).

Analysis of Soluble Phenolics and Amino Acids

Three-month-old, fully grown, still-green plants of the four genotypes (10 wild-type plants, 6 plants of pal1, 6 plants of pal2, and 7 plants of pal1 pal2) were harvested after flowering had ceased. Whole inflorescence stems (without leaves, siliques, or flowers) were immediately frozen and homogenized in liquid nitrogen. Soluble phenolics were analyzed as described by Meyermans et al. (2000). Briefly, the material was extracted with methanol and the supernatant dried and extracted with cyclohexane/water containing 0.1% trifluoroacetic acid (1:1 [v/v]). Phenolics in the aqueous phase were separated and analyzed by reversephase HPLC. Absorbance spectra were recorded with a 996 diode array detector (Waters, Milford, MA) scanning from 200 to 450 nm. The peak height was quantified at the maximum absorbance value between 250 and 450 nm. Data collection and integration were done using the Millenium software (Waters). Peak height/dry weight of 89 peaks (subtracting the major but indifferent peak at retention time of 5.46 in the beginning of the chromatogram) were analyzed with weighted one-way ANOVA, with P = 0.05 for the F test as well as for the least square difference post-hoc test.

Liquid chromatography-mass spectrometry analysis of the water phase was performed using a gradient separation from 83% solvent A (aqueous, 1% acetic acid) to 77% solvent B (acetonitrile, 1% acetic acid) in 21 min on a reverse-phase Luna C18 column (150 \times 2.1 mm, 3 μ m; Phenomenex, Torrance, CA) at a flow of 0.30 mL min⁻¹ (SpectraSystem P1000XR HPLC pump; Thermo Separation Products, Riviera Beach, FL) and a column temperature of 40°C (SpectraSystem AS1000 autosampler; Thermo Separation Products). UV/VIS absorption spectra were taken between 200 and 450 nm on a SpectraSystem UV6000LP detector (Thermo Separation Products). Full MS spectra were obtained in the positive or negative ionization mode on a LCQ Classic MS instrument (ThermoQuest, San Jose, CA), using atmospheric pressure chemical ionization as ion source. Vaporizer temperature, capillary temperature, sheath gas, aux gas, and the source current were set at 450°C, 150°C, 64, 3, and 5 µA, or at 400°C, 150°C, 23, 6, and 5 µA in the positive or negative ionization mode, respectively. MSn spectra were obtained by single reaction monitoring using a collision energy of 35%.

The methanol extracts were also used for amino acid analysis. Of the extracts, 5 μ L was directly applied to an amino acid analyzer (420A derivatizer with an online 130A separation system; Applied Biosystems, Foster City, CA). Amino acids were identified by cochromatography with authentic standards. Peak height/dry weight were analyzed with one-way ANOVA, with P = 0.05 for the *F* test and the Dunnett's post-hoc test.

The lyophilized scopolin peak and synthetic scopolin were subjected to acid hydrolysis (1 N HCl, 3 h at 90°C). The reaction products were resolved on HPLC, as described above, and authenticated with standards of scopoletin and isoscopoletin. Synthetic compounds for authentication were vanillate (Acros, Geel, Belgium), scopoletin (Sigma-Aldrich, St. Louis, MO), isoscopoletin (Fluka, Buchs, Switzerland), and ${\bf FA}(4\text{-}O\text{-}8){\bf G}$ (Ralph et al., 1992). ${\bf FA}(8\text{-}5){\bf G}$ was isolated from the mixture obtained by cross-coupling coniferyl alcohol with ethyl FA via MnO2 in

acetone followed by saponification (H. Kim and J. Ralph, unpublished data). Scopolin was synthesized from scopoletin (Chaudhury et al., 1948).

Substrate Feeding Assays and Thin-Layer Chromatography of Sinapate Esters

Substrate feeding assays with 1 mM of cinnamic acid or *para*-coumaric acid were performed on 15 cm inflorescence stems. Stems were cut under water to avoid collapse of the vascular stream and placed into water with the respective substrate for 24 h. Stem pieces from above the part that was in contact with the substrate were harvested, immediately frozen, and analyzed as above. Thin-layer chromatography of sinapate esters were run as described by Lehfeldt et al. (2000) using leaves and inflorescence stems of 10-week-old greenhouse-grown plants.

Acetyl Bromide Soluble Lignin Determination and DFRC Analysis of Lignin

The inflorescence stems of five plants of each C24, pal1, pal2, and pal1 pal2 were harvested individually after 3 months of growth. Plants were green, and flowering had ceased. Stem pieces were immediately frozen in liquid nitrogen and freeze-dried. The ABSL assay was performed as detailed (Fukushima and Hatfield, 2001). ABSL was calculated using an extinction coefficient of 17.2. DFRC lignin analysis was performed essentially as described previously (Lu and Ralph, 1997; Ralph and Lu, 1998). Monomer determination was performed using response factors (coniferyl diacetate (G), 1.39; sinapyl diacetate (S), 1.44) derived from monomer standards against the internal standard (4,4'-ethylidenebis-phenol; Aldrich, Milwaukee, WI).

Cell Wall Isolation and Saponification

Inflorescence stem material (1.5 g) of the wild type and double mutant was ground in liquid nitrogen and taken up in 30 mL 50 mM NaCl. This mixture was kept at 4°C overnight. Samples were centrifuged for 10 min at 3500 rpm, and the pellet was extracted with 40 mL 80% ethanol and sonicated for 20 min. This extraction was repeated twice. Centrifugation, extraction, and sonication were repeated exactly with the following extractives: acetone, chloroform:methanol (1:1), and acetone. Samples were allowed to air dry. Saponification of isolated cell walls was performed overnight in 2 M NaOH at room temperature under nitrogen. Extracts were neutralized with 2 M HCl under nitrogen and centrifuged. The supernatant was analyzed with liquid chromatography–mass spectrometry as above.

Carbohydrate Analysis

The xylem tissue from the stems was dried and ground (100 mesh). The powder was extracted successively with a mixture of toluene and ethanol (2:1 [v/v]), with ethanol, and with hot water, and then freeze-dried. The cell wall material was hydrolyzed by the $72\%\ H_2SO_4$ method and the sugars analyzed by gas chromatography of the alditol acetate derivatives as described (Chambat et al., 1997).

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